

# Multiplex core-periphery organization of the human connectome

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## Abstract

What is the core of the human brain is a fundamental question that has been mainly addressed by studying the anatomical connections between differently specialized areas, thus neglecting the possible contributions from their functional interactions. While many methods are available to identify the core of a network when connections between nodes are all of the same type, a principled approach to define the core when multiple types of connectivity are allowed is still lacking. Here we introduce a general framework to define and extract the core-periphery structure of multi-layer networks by explicitly taking into account the connectivity patterns at each layer. We first validate our algorithm on synthetic networks of different size and density, and with tunable overlap between the cores at different layers. We then use our method to merge information from structural and functional brain networks, obtaining in this way an integrated description of the core of the human connectome. Results confirm the role of the main known cortical and subcortical hubs, but also suggest the presence of new areas in the sensori-motor cortex that are crucial for the intrinsic brain functioning. Taken together these findings provide fresh evidence on a fundamental question in modern neuroscience and offer new opportunities to explore the mesoscale properties of multimodal brain networks.

**Keywords:** Complex networks, Multilayer networks, Core-periphery structure, Rich-club, Brain connectivity, Multimodal integration

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## 1. Introduction

Complex networks are characterized by the existence of non-random structures at different topological scales [5–7]. A peculiar structure is the so-called core-periphery organization [8], where the network exhibits a group of tightly connected nodes (i.e. the *core*), and a group made by the remaining weakly connected nodes (i.e. the *periphery*).

Core-periphery organization has been recognized as a fundamental property of complex networks to support integration of information [9–16]. A related concept is that of rich-club behavior, where the tightly connected nodes are the network hubs, i.e. the nodes with a large number of links [17, 18]. A rich-club organization has been observed in various real-world systems, such as social, technological and biological networks [17–20], including the brain [21–24]. More recently, a refined version of the rich-club analysis, based not only on the number of connections of the hubs, but also on their capability to bridge different communities, has been shown to be relevant to support the integrative properties of the nervous system [25].

In the human brain, rich-club and rich-core organization, associated to the efficiency in communication and distribution of information, have been mainly reported in anatomical, or structural, connectivity networks obtained experimentally from diffusion tensor imaging (DTI) data. It has been conjectured that rich cores, rather than shortest paths, may actually be responsible for the efficient integration of information between remote brain areas [21], which is a crucial prerequisite for normal cognitive performance [26, 27]. Current evidence suggests that posterior medial and parietal cortical regions mainly constitute the core of the human connectome [21, 28], while they are contradictory on the role of other areas, such as the medial prefrontal cortex (mPFC) and the sensori-motor system, which are basic components of the brain functioning [49]. Because brain regions are also characterized by functional interactions inferred from neuroimaging data, such as functional magnetic resonance imaging (fMRI) [29, 30], we hypothesize that integrating information from both structural and functional networks can give a more accurate estimate of the regions that eventually constitute the core of the human cortex.

Instead of aggregating the two different types of connectivity or analyzing them separately, we adopt a multiplex network approach that preserves and exploits the original information on how brain regions are structurally and functionally interconnected. In a multiplex network, different connectivity types are mathematically represented as networks at different layers. Notably, in a multiplex - a particular case of multilayer network - there is a one-to-one correspondence between the nodes at different layers [31–35]. Multiplex network theory has been recently used to successfully extract higher-order properties of multimodal [36] and multifrequency brain networks that cannot be retrieved by standard approaches [37, 38].

Interestingly, the detection of core-periphery organization in multiplex networks has been poorly explored, with the exception of approaches based on k-core decomposition [42, 43]. To address this gap, we introduce a criterion to define and detect core-periphery organization in multiplex networks. Our method works for any number of layers and is scalable to large networks, being non-parametric and based on local node information [20]. In the following, we first introduce the general framework and then we validate it on synthetic multiplex networks with tunable core similarity.

We finally apply our method to integrate information from structural and functional brain networks and extract the multiplex core-periphery organization of the human brain.

The obtained results point to the main hubs known in the literature, but also allows to highlight the central role played by the regions of the sensori-motor system, which has been surprisingly neglected by previous studies on core-periphery organization, despite being considered a fundamental component of the default-mode network [49]. Our research shades new light on the emergence of core regions in the human connectome, and we hope it will spur further work towards a better understanding of the complex relationships in the nervous system.

## 2. Results

### 2.1. Extracting the rich core of a multiplex network

Let us consider a multiplex network described by a vector of adjacency matrices  $\mathcal{M} = \{A^{[1]}, \dots, A^{[M]}\}$ , where all interactions of type  $\alpha$ ,  $\alpha = 1, \dots, M$ , are encoded in a different layer described by the adjacency matrix  $A^{[\alpha]}$ . To detect the core-periphery structure of a multiplex network, we first compute the multiplex degree vector  $\mathbf{k}_i = \{k_i^{[1]}, \dots, k_i^{[M]}\}$  of each node  $i$  [34]. From now on, we refer to  $k_i^{[\alpha]}$ ,  $\alpha = 1, \dots, M$ , as the *richness* of node  $i$  at layer  $\alpha$ . Notice that this is the simplest way to define the richness of a node, and different measures of richness, such as other measures of node centrality, can be as well used.

For each layer  $\alpha$ , we then divide the links of node  $i$  in those towards lower richness nodes, and those towards higher richness nodes, so that we can decompose the degree of node  $i$  at layer  $\alpha$  as  $k_i^{[\alpha]} = k_i^{[\alpha]-} + k_i^{[\alpha]+}$ . Finally, the *multiplex richness*  $\mu_i$  of node  $i$  is obtained by aggregating single-layer information:

$$\mu_i = \sum_{\alpha=1}^M c^{[\alpha]} k_i^{[\alpha]}. \quad (1)$$

where the coefficients  $c^{[\alpha]}$  modulate the relative relevance of each layer and can, for instance, be determined by exogenous information. In analogy to the single-layer case, we define the multiplex richness of a node towards richer nodes as:

$$\mu_i^+ = \sum_{\alpha=1}^M c^{[\alpha]} k_i^{[\alpha]+}. \quad (2)$$

In the most simple set-up we can assume  $c^{[\alpha]} = c = 1/M \forall \alpha$ . More general functional forms to aggregate the contributions from different layers, giving rise to alternative measures of  $\mu_i$  and  $\mu_i^+$ , are presented in the Methods section.

The nodes of the multiplex are ranked according to their richness  $\mu$ , so that the node  $i$  with the best rank, i.e.  $rank_i = 1$ , is the node with the largest value of  $\mu$ , the node ranked 2 is the one with the second largest value of  $\mu$ , and so on. We then plot for each node  $i$  the value of  $\mu_i^+$  as a function of  $rank_i$ . The value of the rank corresponding to the maximum of  $\mu_i^+$  finally determines the core-periphery structure. All nodes with rank lower than such a value are assigned to the multiplex core, whereas the remaining ones become part of the periphery. We notice that also in the simplest case, when  $c^{[\alpha]} = c \forall \alpha$ , the multiplex core-periphery partition cannot be obtained by simply combining the cores

of the different layers, or by applying the single-layer algorithm on the corresponding aggregated network.

As an illustrative example, we report in Fig. 1 the curve  $\mu_i^+$  as a function of  $rank_i$  obtained in the case of the Top Noordin Terrorist network, a multiplex network of  $N = 78$  individuals with three layers (encoding information about mutual trust, common operations and exchanged communication between terrorists), which has been used as a benchmark to test measures and models of multiplex networks [34].

Coefficients  $c^{[\alpha]}$  were chosen, in this case, to be inversely proportional to  $K^{[\alpha]}$  to compensate for the different densities of the three layers. The resulting multiplex rich core integrates information from all the layers and looks different from the rich cores obtained at each of the three layers by a standard single-layer rich core analysis. More details about the results of this analysis are reported in Table S1.

## 2.2. Testing the method on multiplex networks with tunable core similarity

A network with a well defined core-periphery structure has a high density of links among core nodes. With a suitable labeling of the nodes, the adjacency matrix of the network can be decomposed into four different blocks: a dense diagonal block encoding information on core-core links, a sparser diagonal block describing links among peripheral nodes, and two off-diagonal blocks encoding core-periphery edges. The key feature of such block-structure is that  $\rho_1 \gg \rho_3$ , i.e. the density  $\rho_1$  of the core-core block is much higher than that of the periphery-periphery block,  $\rho_3$ . As first noted by Borgatti and Everett [8], the density  $\rho_2$  of the off-diagonal blocks is typically not a crucial factor to characterize a core-periphery structure.

To test how our method works on multiplex networks with different structures, we have introduced a model to produce synthetic multiplex networks with tunable core similarity. In particular, we have constructed multiplexes where each of the  $M = 2$  layers contain  $N = 250$  nodes and only  $N_c = 50$  of them belong to the core. Each layer has the same average node degree  $\langle k \rangle = 10$ , and the same set of parameters  $\rho_1 > \rho_2 > \rho_3$  to describe its core-periphery structure. Our model allows to control the number of nodes that are both in the core of layer 1 and 2. (see Methods for more details).

To quantify the similarity among cores at different layers, we introduce the core similarity  $S_c^{[\alpha]}$  of layer  $\alpha$  with respect to the other layers as:

$$S_c^{[\alpha]} = \frac{1}{(M-1)} \sum_{\beta \neq \alpha}^M \frac{I_c^{[\alpha\beta]}}{N_c^{[\alpha]}}, \quad (3)$$

where  $I_c^{[\alpha\beta]}$  is the number of nodes in the core of both layer  $\alpha$  and layer  $\beta$ , whereas  $N_c^{[\alpha]}$  is the size of the core at layer  $\alpha$ . The core similarity  $S_c^{[\alpha]}$  ranges in  $[0, 1]$ . When layer  $\alpha$  does not share core nodes with any other layers we have  $S_c^{[\alpha]} = 0$ , when all its core nodes also belong to the cores of the other layers  $S_c^{[\alpha]} = 1$ , and when on average only half of them are part of the cores on each other level  $S_c^{[\alpha]} = 1/2$ . The average core similarity of the multiplex can then be computed as  $S_c = (1/M) \sum_{\alpha=1}^M S_c^{[\alpha]}$ .

In Fig. 2 we show the results for three multiplex networks with different core similarity. In the left column of Fig. 2 we consider a multiplex with  $S_c = 0$ . The cores of the two layers are not overlapping, as shown in panel (a). As a consequence, many nodes with high degree in one layer have low degree in the other one. When  $c^{[1]} = c^{[2]} = 0.5$ , the

126 multiplex core of the system is formed by those nodes with sufficiently high multiplex  
 127 richness, as shown in panel (b). In panel (c) we show the changes in the multiplex core  
 128 when we partially ( $c^{[1]} = 0.75, c^{[2]} = 0.25$ , left subplot) or completely ( $c^{[1]} = 1, c^{[2]} = 0$ ,  
 129 right subplot) bias the algorithm towards the first layer.

130 In the central column of Fig. 2 we consider a multiplex with  $S_c = 1/2$ . Half of the  
 131 core nodes are in common to both layers while half are typical of each layer. The block  
 132 structure of the two layers is partially overlapping, and the nodes are spread uniformly  
 133 over the  $k_i^{[2]}$  vs  $k_i^{[1]}$  plane. In the unbiased case the multiplex core of the system is  
 134 formed by nodes which are part of the core on both layers, but also by nodes scoring  
 135 extremely high in one layer, despite being in the periphery in the other one (panel b).  
 136 When  $c^{[1]} > c^{[2]}$ , this is particularly true for nodes which have high richness in the first  
 137 layer and low richness in the second, while the opposite is much more unlikely (panel c).  
 138 In the right column of Fig. 2 we consider a multiplex with  $S_c \approx 1$ . The block structure  
 139 of the two layers is now almost identical; the node degrees  $k^{[1]}$  and  $k^{[2]}$  are correlated  
 140 and most of the nodes belonging to each core are in the multiplex core (panel b). As  
 141 the core structure at the two layers are extremely similar, the biased cases do not differ  
 142 significantly from the unbiased one (panel c).

### 143 2.3. Merging structure and function to extract the connectome's core

144 We have applied our method to investigate the human connectome by considering, at  
 145 the same time, structural and functional information. We have therefore constructed a  
 146 multiplex brain network formed by one structural layer and one functional layer. The two  
 147 layers were obtained by first averaging brain connectivity matrices estimated respectively  
 148 from DTI and fMRI data in 171 healthy individuals. Each of the two layers is then  
 149 thresholded by fixing the average node degree  $\langle k \rangle$ . We have focused our analysis on 158  
 150 regions of interest (ROIs) of the cortex (see Methods for more details).

151 In Fig. 3 we report the cores found by analyzing separately the two layers, as well  
 152 as the multiplex core obtained with our method. The figure refers to the case of a  
 153 representative threshold corresponding to an average node degree  $\langle k \rangle = 7$ . We notice  
 154 that the cores of the structural and functional layers are only partially overlapping, with  
 155 a value of core similarity of  $S_c = 0.15$ . For the sake of completeness, we also report  
 156 the  $S_c$  values for the the entire threshold range (Fig. S1). **A detailed analysis on the**  
 157 **robustness of the multiplex core detection in presence of random fluctuations is reported**  
 158 **in the Supplementary text S1.**

159 As shown in Fig. 3, ventral brain areas tend in general to form the structural core,  
 160 while more dorsal regions appear in the functional core. Notably, brain regions of interest  
 161 (ROIs, Table S2) that are in the core of both structural and functional layers also tend to  
 162 be in the core of the multiplex. Instead, ROIs being in the periphery of both layers tend  
 163 to be excluded from the multiplex core. However, exceptions may exist depending on the  
 164 multiplex richness of the nodes. For example, the posterior part of the right precentral  
 165 gyrus (RCGa3), which is in the periphery of both the structural and functional layer,  
 166 is eventually assigned to the multiplex core, because of its relatively high rank score in  
 167 the two layers. The situation appears even less predictable for ROIs that are in the  
 168 core of one layer and in the periphery of the other layer. Only occasionally these will  
 169 belong to the multiplex core. This is the case, for example, of the anterior part of right  
 170 precentral gyrus (RCGa2) which exhibits a relatively low structural richness but high

functional richness, i.e. ranked seventh in the functional core, or of the anterior part of the right parietal operculum (RPOC1), which has the highest structural richness but a low functional richness.

#### 2.4. *Revealing new core regions of the human brain*

We have extracted the multiplex core-periphery structure of the human brain for the full range of available thresholds  $\langle k \rangle = 1, 2, \dots, 120$  (see Methods for more details). In this way, we have been able to calculate the *coreness*  $C_i$  of each node  $i$ , defined as the normalized number of thresholds at which the corresponding ROI is present in the rich core. This allows us to rank ROIs according to their likelihood to be part of the multiplex core and to compare these to the rankings obtained separately for structural and functional layers. We note that the same approach of investigating the persistence across a set of different filtering thresholds can be applied to any node property. This can turn useful for statistical validation in the case no threshold is universally accepted, as often happens for brain networks [44–46].

Parietal (pre/cuneus PCU/LOC, superior parietal lobe SPL), cingulate (anterior Ca, posterior Cp), temporal (superior temporal gyrus), insular (insular cortex IC), as well as frontal ROIs (paracingulate PC) mainly constitute the structural core, as shown in Fig. S2. While some overlap exists between the structural and the functional cores, the latter rather tends instead to include occipital (occipital fusiform gyrus OFG, temporo-occipital fusiform cortex TOFC) and central (pre/post central gyrus CGa/CGp) ROIs and, notably, to exclude regions in the frontal lobe (top 25% ROIs, Fig. S3).

Fig. 4 shows the coreness of the multiplex network. As expected, ROIs that are peripheral (i.e., low coreness) in both layers are also peripheral in the multiplex, while ROIs with both a high structural and high functional coreness are typically observed in the multiplex core (e.g., TOFC, OFG, Ca, Cp). Interesting behaviors emerge for those regions typically characterized by high coreness in one layer and low coreness in the other layer. In fact, some of these ROIs are part of the multiplex core, while others are usually found in the multiplex periphery, as shown Fig. 5a. For areas with a different assignment in the two layers, we note that the main contribution to the multiplex richness  $\mu_i$  comes from the richness in the layer where node  $i$  is identified as core. Interestingly, not only the average richness of the node in the core layer is higher than the one in the peripheral layer, but also its fluctuations around the mean.

As a consequence, among regions that are core in the structural layer but peripheral in the functional one, those with relatively higher structural richness (degree), such as precuneus PCU, insular cortex IC and posterior cingulate Cp, finally tend to join the multiplex core no matter the exact value of their functional richness (upper right corner of Fig. 5a). Conversely, ROIs with relatively lower structural degree are usually peripheral in the multiplex, and typically located in the pre-frontal cortex PC and frontal lobe FP (lower right corner of Fig. 5a), as illustrated in Fig. 5b,c. Similarly, among areas in the functional core, those with relatively higher functional degree, such as precentral gyrus CGa and central operculum COC, tend to join the multiplex core (upper left corner of Fig. 5a). In contrast, ROIs with relatively lower functional degree, are mostly peripheral in the multiplex, and are located in the parietal operculum POC and superior frontal gyrus SFG (lower left corner of Fig. 5a).

In a separate analysis, we have extracted the multiplex brain coreness from each individual and we show that, despite a normal inter-subject variability, the average mul-

tiplex brain coreness is very similar to the multiplex coreness of the group-averaged brain networks (Fig. S4). Finally, we have evaluated the robustness of the results when also including subcortical ROIs in the brain networks. We report that thalamus, putamen and hippocampus are among the regions with highest coreness and therefore become part of the multiplex core (Fig. S5). Interestingly, their presence does not significantly alter the coreness of the other ROIs (Fig. S6), suggesting an assortative structure where highly connected subcortical regions preferentially get connected with regions in the cortex.

### 3. Discussion

The existence of a network core in the brain is a prerequisite for neural functioning and cognition, and damages to the core have been associated with several neurological or psychiatric diseases [23, 47, 48]. Finding the router regions that ensure integration between the different brain modules and communication in the system is therefore a fundamental question in neuroscience. Previous studies have addressed the question by considering only the structural connectivity of the brain through disparate techniques, such as  $k$ -core decomposition, centrality measures, and rich-club analysis [21, 28]. While the obtained results agree on the implication of posterior medial and parietal cortical regions - as well as subcortical thalamus, putamen and hippocampus - in the network core [21, 28], they neglect the possible role of other areas which are crucial from a functional perspective, such as those in the default-mode network (DMN) [49].

To integrate information from both structural and functional brain connectivity at the network level, we introduce a general criterion to define and extract the core when nodes are connected through links which can vary in meaning and nature, and the whole system can be described as a network with multiple layers [31–35]. Compared to standard approaches, this method has the theoretical advantage to provide a more robust solution, taking into account the relative importance of the nodes at each layer, rather than simply considering the union or intersection of the cores across layers, or extracting the core from the aggregated network.

The obtained results shed new light on the role of the regions characterizing the intrinsic brain function to eventually shape the core of the human brain. First, we show that mPFC (e.g., PC and FP), exhibiting a high structural but low functional coreness, is eventually assigned to the periphery (Fig. 5a, lower-right corner). This outcome can be predicted by the lower multiplex richness and relatively low structural degree, and not by the solely attitude of frontal areas to be peripheral in the functional brain network (Fig. 5b,c). The exclusion of the mPFC from the rich core supports the hypothesis that default-mode network activity may be mainly driven from highly coupled areas of the posterior medial and parietal cortex, which in turn link to other highly connected regions, such as the medial orbitofrontal cortex [28].

Second, while frontal ROIs are excluded, new regions gain importance and become part of the core because of their higher multiplex richness (see Fig. 5a, upper left corner). Among them, we report areas of the central gyrus (CGa, CGp to a minor extent), which are characterized by a low structural but relatively high functional degree, as shown in Fig. 5b,c. These regions are part of the primary sensori-motor cortex, which has been shown to be the most extensive of the resting-state components, or networks (out of 8 [50]), covering 27% percent of the total gray matter in the brain [51]. The primary sensori-motor component has a high degree of integration (overlap and activity



coupling) with all other resting-state networks (e.g., DMN), which is consistent with the increased synchronization of neural activity in cortical regions during sensory processing [52]. Notably, ongoing functional connectivity in the primary sensori-motor network, originally revealed by seed-based analysis [53, 54], has been extensively verified by ICA and clustering methods [55, 56].

Our method provides an effective tool to integrate mesoscale topological information in brain networks derived from multimodal neuroimaging data. Multimodal integration of brain networks is gaining more and more interest [57–60] due, on the one hand, to the increasing availability of large heterogenous datasets (e.g. HCP <http://www.humanconnectomeproject.org>, ADNI <http://adni.loni.usc.edu>) and, on the other hand, to the need of principled ways to characterize multiscale neural mechanisms (e.g., cross-frequency coupling) and provide predictive diagnostics for multifactor brain diseases, such as Alzheimer’s disease.

It is important to note, that our analysis of the human connectome relies on the assumption that each layer contributes with the same intensity to the definition of the multiplex core. In general, however, the contribution of a layer  $\alpha$  can be weighted differently through an opportune choice of the parameter  $c^{[\alpha]}$ , and this can be used to enhance or reduce the importance of the different types of connectivity. A larger value of  $c^{[\alpha]}$  increases the relevance of the corresponding layer until when, in the limit in which  $c^{[\alpha]} \rightarrow 1$  and the coefficients of all the other layers go to zero, the multiplex core is not any more defined by the topology of all the  $M$  layers, but coincides with the core at layer  $\alpha$ . For instance, setting  $c^{[structural]} = 1$  and  $c^{[functional]} = 0$  returns a core based on the anatomical information only, and in agreement with most of the previous literature on such topic (see Fig. S2). As an unbiased way to characterize the multiplex core of the human brain, we have focused our analysis on the simplest and symmetric case,  $c^{[structural]} = c^{[functional]} = 0.5$ . We show in Fig. S7 that the results are relatively stable for small perturbations around this unbiased condition. However, other combinations are in general possible and should be adopted if supported by a plausible rationale. For example, in the case of multifrequency brain networks, one could assign stronger weights to higher frequency layers in order to compensate for  $1/f$  frequency scaling of power spectra [61].

From an operative point of view, the proposed method to detect the core-periphery organization in multiplex networks has two clear advantages: *i*) it is fast and scalable, since it works using only local information; *ii*) it is non-parametric, e.g. no need to input *a-priori* information such as the core size. Moreover, it can be generalized in a straightforward way to the case of directed networks. A drawback of the method is that it focuses on highly connected rich nodes, and neglects the possible important role of the so-called *connectors*, i.e. central nodes with low degree [62]. We note that alternative core-periphery structures which include connectors can be detected by more computationally demanding methods such as those based on stochastic block models, which have been recently proposed to extract the mesoscale structure of time-varying and multilayer networks [63]. We hope that our work can trigger further developments in the exploration of core-periphery structure of real-world large-scale multiplex networks.

To conclude, our method to investigate multiplex core-periphery organization in complex networks suggests that the core of the human cortex is made up of known cortical and subcortical hubs, as well as of areas in the sensori-motor system that were previously overlooked by standard approaches, but that are crucial for the brain functioning. Our



findings offer an augmented definition of the rich core of the human brain, which takes into account not only the anatomical structure but also its function.

We hope that our work will contribute to advance our understanding of the mesoscale connectivity mechanisms in multiplex brain networks, in an effort to better integrate the one-to-many relationships that exist between structure and function in the human brain [29].

## 4. Methods

### 4.1. Multiplex stochastic block model with tunable core similarity

Stochastic block models for multiplex networks have been recently introduced by Peixoto [63]. Here, we introduce a stochastic block model that allows to sample multiplex networks with an assigned value of core similarity  $S_C$  (see Eq. 3). Suppose we have  $N$  nodes and we want to construct a multiplex network having a core-periphery structure at each layer  $\alpha = 1, \dots, M$ , with  $N_c^{[\alpha]}$  nodes in the core of layer  $\alpha$ .

In particular, we set  $M = 2$ ,  $N = 250$ ,  $N_c^{[1]} = N_c^{[2]} = N_c = 50$ , and we create at each layer a core-periphery structure with the same set of densities:  $\rho_1 = 0.2$ ,  $\rho_2 = 0.04$  and  $\rho_3 = 0.03$ . Namely, for each of the two layers, we connect with a probability  $\rho_1$  two nodes both in the core, with probability  $\rho_2$  a node in the core and a node in the periphery, and finally with probability  $\rho_3$  two peripheral nodes. The values of the three parameters were chosen in a way that  $\langle k \rangle = 10$  on both layers, and the core-periphery structure of each layer is sufficiently strong to be detected with good accuracy, as discussed in the Supplementary text S2.

Different levels of core similarity are achieved by varying the overlap between core nodes at the two layers. When the two sets of core nodes are completely overlapping,  $S_c = 1$ , whereas when the two sets are disjoint  $S_c = 0$ . Despite other related formulations of  $S_c$  are possible, our definition reflects the intuition that when two layers with equal core size share half of the core nodes, then  $S_c = 1/2$ .

### 4.2. Multiplex richness $\mu_i$ and $\mu_i^+$

The multiplex richness  $\mu_i$  and  $\mu_i^+$  introduced in Eqs. 1 and 2 are obtained by mean of a simple aggregation of information based on the single layers. In the simplest set-up  $c^{[\alpha]} = c = 1/M$  for  $\alpha = 1, \dots, M$ , and the multiplex richness  $\mu_i$  of a node  $i$  is simply proportional to its overlapping degree  $o_i$  [34]. A layer with higher density weighs more in the computation of the multiplex core of a network.

In general, coefficients  $c^{[\alpha]}$  can be used to modulate the relevance to the layers of the network in order to extract its core. If one wants to have equal contributions to  $\mu_i$  and  $\mu_i^+$  from all the layers but their number of links  $K^{[\alpha]}$  is different - for instance because in some layers it might be easier to establish or measure a connection than in others - a natural choice is to set  $c^{[\alpha]}$  to be proportional to  $1/K^{[\alpha]}$ . In other cases, independently from their density, it might be reasonable to assign different importance to different layers, because of exogenous information. Once again this can be achieved by assigning different values of the coefficients  $c^{[\alpha]}$ .

At last, we notice that Eq. 1 is a particular choice of a more general scenario, where the multiplex richness  $\mu_i$  is a generic function  $f$  of the degree of a node at the different layers:

$$\mu_i = f(k_i^{[1]}, \dots, k_i^{[M]}). \quad (4)$$

and  $\mu_i^+$  is a function of a generic function  $g$ :

$$\mu_i^+ = g(k_i^{+[1]}, \dots, k_i^{+[M]}). \quad (5)$$

#### 4.3. Multimodal brain networks

We have considered 171 healthy human subjects from the NKI Rockland dataset [http://fcon\\_1000.projects.nitrc.org/indi/pro/nki.html](http://fcon_1000.projects.nitrc.org/indi/pro/nki.html). We have used diffusion weighted magnetic resonance imaging (dwMRI) and functional magnetic resonance imaging (fMRI) to derive respectively structural and functional brain networks in each subject.

We have gathered the corresponding connectivity matrices from the USC Multimodal Connectivity Database (<http://umcd.humanconnectomeproject.org>) [64].

In particular, structural connectivity have been obtained using anatomical fiber assignment through the continuous tracking (FACT) algorithm [65]. Functional connectivity has been computed by means of the Pearson's correlation coefficient between fMRI signals recorded during a 10 minute resting state (RS). RS-based functional connectivity measures the amount of interaction - or temporal dependence - between different brain areas during spontaneous brain activity [30]. More details about the processing steps can be found here [66]. A total number of  $N = 188$  regions of interest (ROIs) are available for both structural and functional brain networks, thus resulting in connectivity matrices of size  $N \times N$ , spatially matched with the MNI152 template [67].

Because we are mainly interested in cortical networks, we focused our analysis on the network obtained by removing all subcortical ROIs and obtained connectivity matrices of size  $158 \times 158$ . The results for all the ROIs are reported for the sake of completeness. The full name and acronym for all the ROIs can be found in Table S1. We have then averaged the resulting connectivity matrices (after Fisher transformation) across subjects in order to have a population-level representation. At the end, we have obtained a structural weighted connectivity matrix  $\mathcal{S}$ , whose entry  $s_{ij} = s_{ji}$  contained the group-average number of axonal fibers between ROIs  $i$  and  $j$ , and a functional weighted connectivity matrix  $\mathcal{F}$ , whose entry  $f_{ij} = f_{ji}$  corresponded to the group-average correlation coefficient between the fMRI signals of ROIs  $i$  and  $j$ .

We have used density-based thresholding to derive structural and functional brain networks by removing the lowest values from the connectivity matrices and binarizing the remaining ones [30]. We have considered a full range of density thresholds, corresponding to an increasing average node degree  $\langle k \rangle = 1, 2, \dots, 120$ . The last value was given by the maximal  $\langle k \rangle$  observed in the native structural connectivity matrices, which are originally not fully connected. After filtering, for each threshold we have combined the resulting structural and functional brain networks into a multiplex network  $\mathcal{M} = \{\mathcal{S}, \mathcal{F}\}$ .

#### Competing interests

We have no competing interests.

## Authors' contributions

FB carried out the theoretical work, participated in data analysis, participated in the design of the study and drafted the manuscript; JG participated in data analysis and drafted the manuscript; MC conceived and designed the study, and helped draft the manuscript. VL conceived the study and drafted the manuscript; FDVF coordinated the study, participated in the design of the study, participated in data analysis and drafted the manuscript. All authors gave final approval for publication.

## Funder

FDVF and MC acknowledge support by the ANR French program through the contracts ANR-10-IAIHU-06 and ANR-15-NEUC-0006-02. The funders had no role in study design, data collection and analysis, decision to publish, or preparation of the manuscript.

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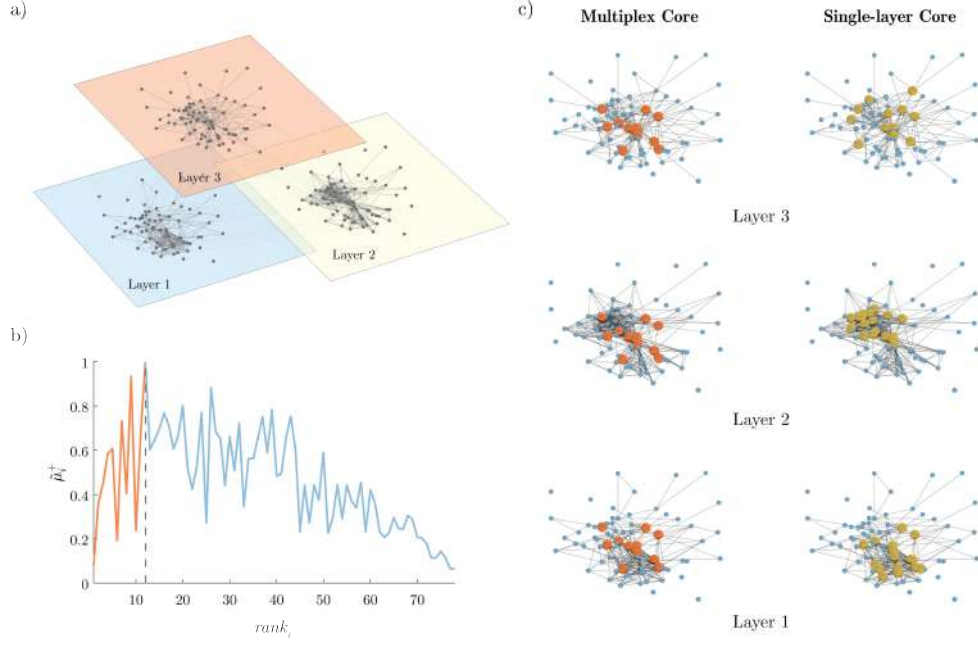
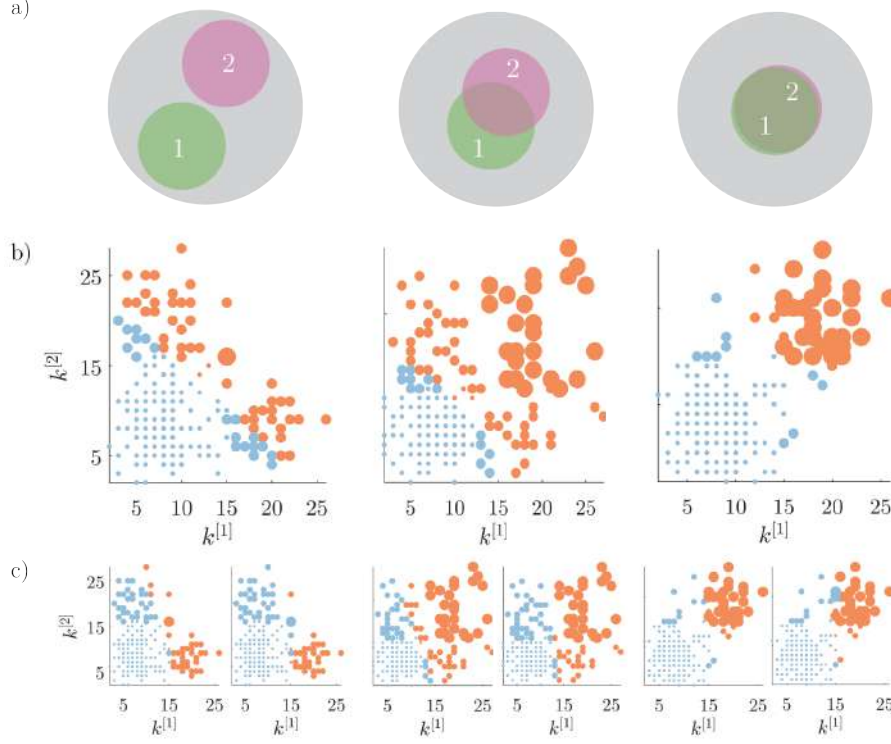
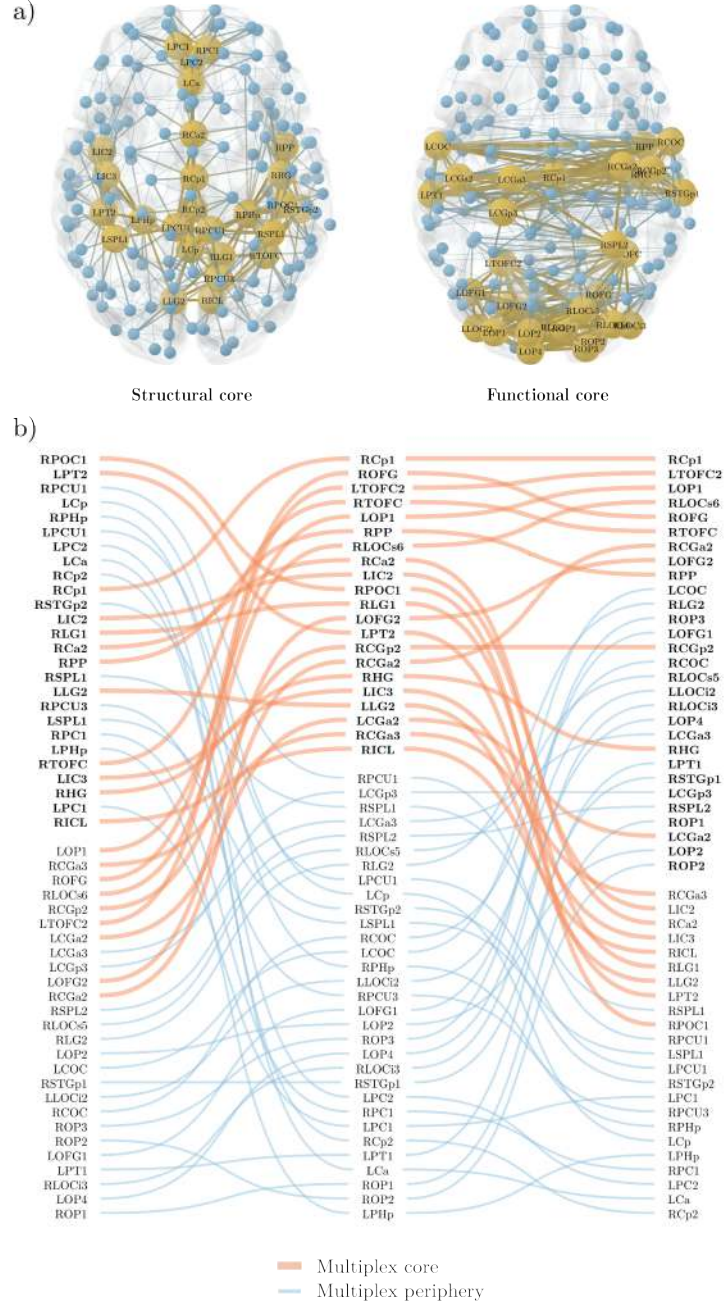


Figure 1: **An illustrative example of the multiplex rich core analysis.** In panel (a) we show a multiplex social network obtained from the Top Noordin Terrorists' contacts, with  $N = 78$  nodes,  $M = 3$  layers and  $K^{[1]} = 259$ ,  $K^{[2]} = 437$  and  $K^{[3]} = 200$ , for the three layers respectively. Panel b) shows the curve  $\tilde{\mu}_i^+ = \mu_i^+ / \max(\mu_i^+)$  as a function of  $rank_i$ . All nodes from rank equal to 1 up to the node with maximum  $\tilde{\mu}^+$  are part of the core of the multiplex, which is shown in red color in panel (c), first column. The cores obtained at each layer by the standard single-layer analysis are reported in yellow for the sake of comparison in the second column. The percentages of core nodes in the single layers that are in the multiplex core are respectively 83.3% for layer 1, 66.7% for layer 2, and 58.3% for layer 3





**Figure 2: Core-periphery structure in synthetic multiplex networks with different core similarity.** In panel (a) we sketch multiplex networks with  $M = 2$  layers,  $N = 250$  nodes and different levels of core similarity, namely  $S_c = 0$  (left column),  $S_c = 1/2$  (central column) and  $S_c = 1$  (right column). In panel (b) the nodes are placed in a two dimensional plane according to their degree at each layer. The size of each dot is proportional to the multiplex richness  $\mu_i$  of the node (with  $c^{[1]} = 1, c^{[2]} = 0$ ). Nodes belonging to the multiplex cores are usually placed in the right-top corner of the plots and are colored orange, while the multiplex periphery is in blue. In panel (c) we report results obtained for two cases with  $c^{[1]} \neq c^{[2]}$ , namely:  $(c^{[1]} = 0.75, c^{[2]} = 0.25)$  where the core is biased towards the important nodes of the first layer (left), and  $(c^{[1]} = 1, c^{[2]} = 0)$ , where the core corresponds to the core of the first layer (right).



**Figure 3: Extracting the multiplex core of the human brain from structural and functional information.** Panel a) The structural and functional brain networks filtered with an average node degree  $\langle k \rangle = 7$  are shown respectively on the left and right side. They are represented from above with the frontal lobe pointing upward. The position of the nodes corresponds to the actual location of the brain regions of interests (ROIs, Table S2). Yellow and large nodes represent the brain regions belonging to the core according to the standard single-layer method. Blue and small nodes code for the ROIs in the periphery. Links are yellow and thick if they connect two ROIs in the core, while they are blue and thin if they connect two peripheral nodes. Panel b) ROIs are ranked from top to bottom according to their richness in the structural (left column), functional (right column) and multiplex network (central column). In each column, the labels in bold/normal font stand for the ROIs that are in the core/periphery. For the sake of simplicity, only ROIs that are at least in one core (structural, functional or multiplex) are listed in the three columns. Red/blue and thick/thin lines identify ROIs that go into the core/periphery according to the multiplex approach.

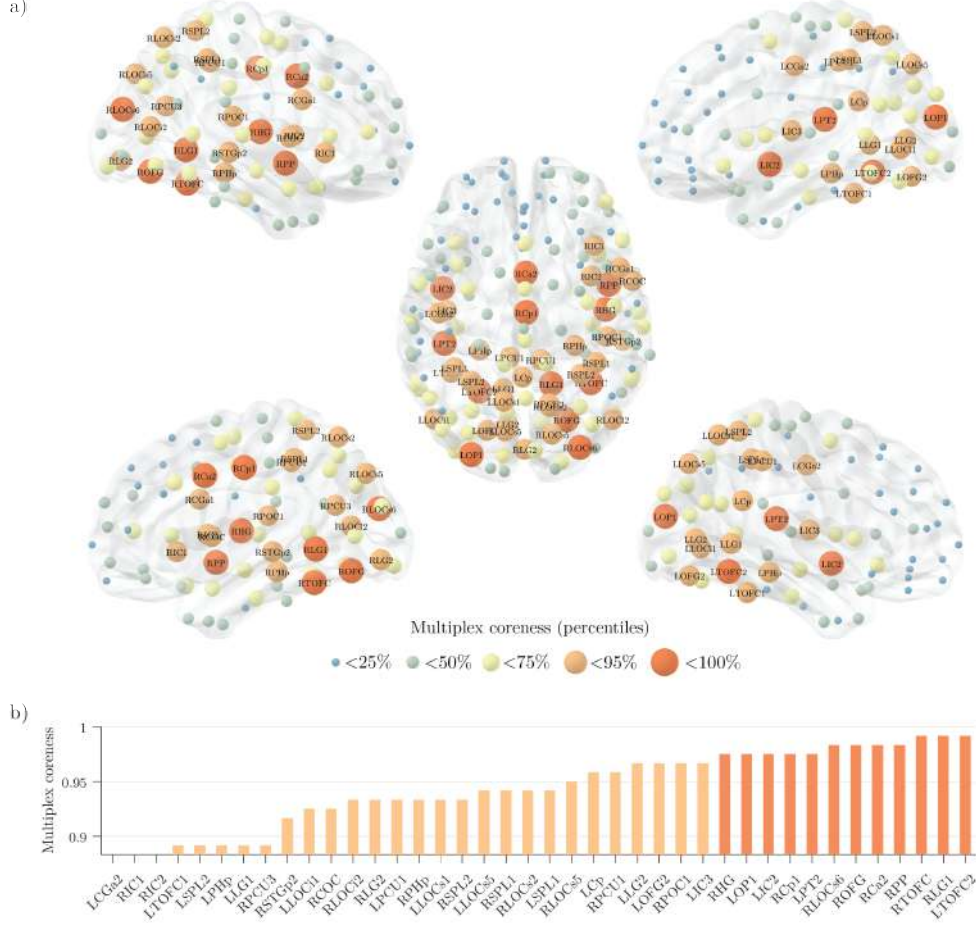


Figure 4: **The multiplex core of the human connectome.** Panel (a) shows the human brain, where regions of interest (ROIs) are highlighted based on their multiplex coreness. The color and size of the nodes are associated to the percentiles of multiplex coreness in each brain region, so that core nodes are larger in size and coloured in red. Left side shows the lateral view of the left hemisphere (top=dorsal, bottom=ventral). Right side shows the lateral view of the right hemisphere (top=dorsal, bottom=ventral). In the middle, the brain is shown from above, with the frontal lobe pointing upward. In panel (b) we report the ROIs corresponding to the 25% highest values of multiplex coreness. The color follows the same legend as in panel (a).

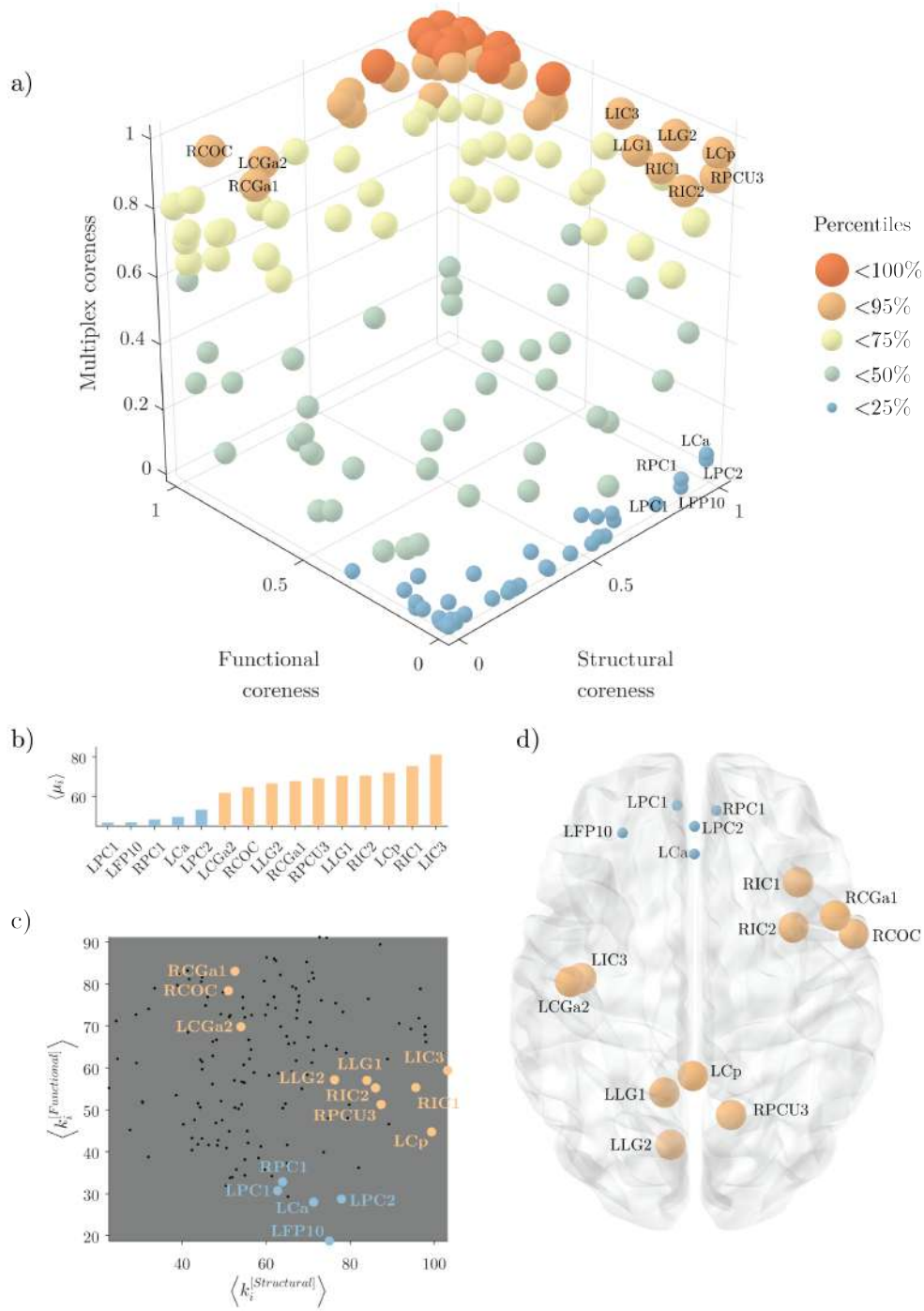


Figure 5: **Emergent non-trivial core regions in the multiplex brain.** Panel (a) shows the scatter plot of the structural, functional and multiplex coreness of the regions of interest (ROIs) in the brain. The color and size of the nodes are associated to the percentile of multiplex coreness across the set of brain regions, as in Fig. 4. Panel (b) reports the average value of multiplex richness  $\langle \mu_i \rangle$  across the different thresholds for the ROIs with the strongest differences in structural and functional coreness. The color follows the same legend as in panel (a). Panel (c) illustrates the distribution of the ROIs (black points) as a function of their averaged structural and functional degree across all the thresholds. Only the ROIs listed in panel (b) are highlighted according to the same color legend as in panel (a).

### S1. Robustness of the rich-core detection

To test the robustness of our method, we have implemented a simulation model allowing random fluctuations in the distributions of values of the multiplex richness  $\mu^+$ . In particular, we have modified the value of each node  $i$  so that the new  $\mu_i^+ = \mu_i^+ + \eta_i \mu_{i,\max}^+$ . Here,  $\mu_{i,\max}^+$  is the maximum value of the original richness and  $\eta_i$  is a random variable within the range  $[-\eta_{\max}, \eta_{\max}]$ , where  $\eta_{\max}$  is a tunable parameter ranging from 0 to 1. Hence, when  $\eta_{\max} = 0$ , the richness of the nodes is not altered; when  $\eta_{\max} = 1$ , the richness of the nodes is independently and maximally altered by a random factor within the range  $[-\mu_{i,\max}^+, \mu_{i,\max}^+]$ . We have applied this simulation model to the multiplex richness values of the brain networks illustrated in the Figure 3 of the main text and we have checked the composition (i.e., the size) of the core as a function of  $\eta_{\max}$ . Notably, for each value of  $\eta_{\max}$  we generate 100 random samples. Results show that the average core size is relatively stable for a broad range of  $\eta_{\max}$  values (Fig. 1a). Notably, fluctuations are completely negligible until  $\eta_{\max} = 0.04$ , and have a high chance to significantly modify the identified rich core only when they are larger than 0.1 (Fig. 1b).

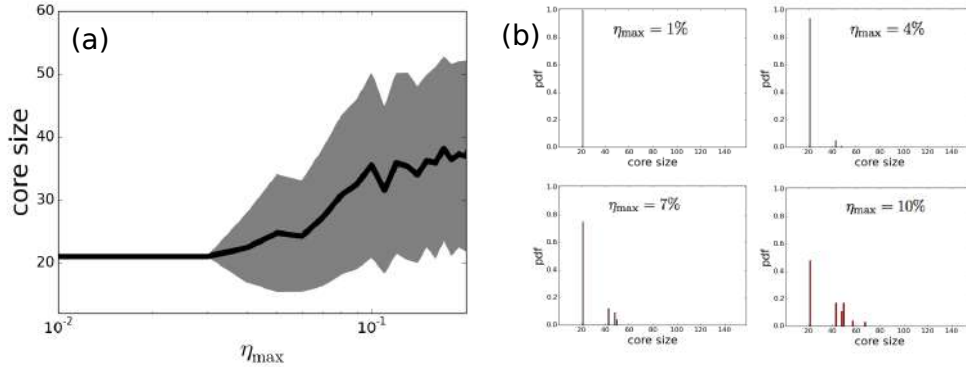


Figure 1: (a) Average core size in the multiplex brain network with  $\langle k \rangle = 7$  as a function of the fluctuations parameter  $\eta_{\max}$ . (b) Cumulative density function of the core size for four selected values in the range  $0.01 \leq \eta_{\max} \leq 0.1$ .

Finally, we have compared the main results with those obtained through an alternative approach where the core-periphery threshold is selected according to a statistical criterion. To this purpose, we have generated 100 degree-constrained random networks from both structural and brain networks. We have then normalized the actual  $\mu^+$  values with respect to those obtained from the random samples according to a standard Z-score  $z(\mu_i^+) = \frac{\mu_i^+ - \bar{\mu}^+}{\sigma(\mu^+)}$ . We eventually report that the regional coreness (on which all the main results are based) is relatively stable regardless whether we have considered the maximum from the actual or normalized values of  $\mu^+$  (Fig. 2).

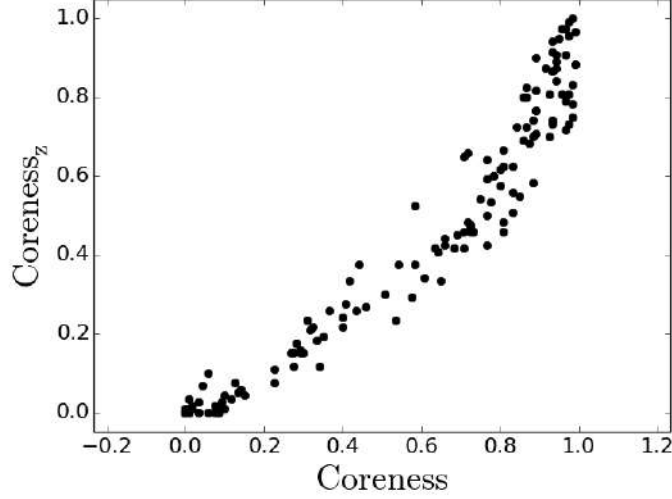


Figure 2: Scatterplot of the multiplex coreness obtained by looking at the maximum values of  $\mu^+$  and that obtained from the maximum of the corresponding Z-scores (Coreness z). The two measures are extremely correlated,  $\rho_s = 0.97, p = 1.04 \times 10^{-80}$ .

## 593 ***S2. Stochastic block model for rich cores in single-layer networks***

594 Suppose we have  $N$  nodes and we want to construct a single-layer network from which  
 595 we can identify a partition into two sets: a core of size  $N_c < N$  and a periphery of size  
 596  $N_p = N - N_c$ . Here we test the performance of the single-layer algorithm to detect rich  
 597 cores [20] on a simple stochastic block model.

598 Let us consider  $N$  nodes from which  $N_c$  drawn at random are chosen to be part  
 599 of the network core, whereas the remaining  $N_p$  are part of the periphery. A network  
 600 with core-periphery structure is such that its adjacency matrix can be decomposed into  
 601 four different blocks: a dense diagonal block encoding information on core-core links,  
 602 a sparser diagonal block describing links among peripheral nodes, and two off-diagonal  
 603 blocks encoding core-periphery edges.

604 In our block model, we connect two nodes with probability  $\rho_1$  if they both belong to  
 605 the core, with probability  $\rho_2$  if one of them belongs to the core and one to the periphery,  
 606 and with probability  $\rho_3$  if they both belong to the periphery,  $\rho_1 \geq \rho_2 \geq \rho_3$ . Given a  
 607 stochastic realization of the block model, we can extract the rich core of the network  
 608 and compare it with the ground-truth, i.e. the set of nodes originally labeled as core  
 609 nodes. In particular, we can test the accuracy of the algorithm for different choice of the  
 610 parameters  $\rho_1$ ,  $\rho_2$  and  $\rho_3$ .

611 Given the three probabilities, the expected total number of edges connecting two core  
 612 nodes is  $K_{cc} = \rho_1[(N_c - 1) * N_c / 2]$ , the expected total number of edges connecting two  
 613 peripheral nodes is  $K_{pp} = \rho_3[(N - N_c - 1) * (N - N_c) / 2]$ , and the expected total number of  
 614 edges connecting a node in the core and a node in the periphery  $K_{cp} = \rho_2[N_c * (N - N_c)]$ .  
 615 The total number of links is  $K = K_{cc} + K_{cp} + K_{pp}$ .



In the case  $\rho_1 = \rho_2 = \rho_3 = \rho$  the nodes are statistically indistinguishable from a structural point of view, the network lacks a core-periphery structure and specifying the value of  $\rho$  simply sets the expected average degree of the network  $\langle k \rangle = N\rho$ . For instance, for  $N = 250$  and  $\rho = 0.04$  we obtain  $\langle k \rangle = 10$  and  $K = 1250$ . Of the different blocks of the adjacency matrix, the exact value of the density of the block encoding links between core and periphery nodes does not play a significant role [8]. For such a reason here we set  $\rho_2 = 0.04$ , and study the core-periphery structure of the network as a function of  $\rho_1$ , with  $\rho_1 > \rho_2$ . The higher the value of  $\rho_1$ , the stronger the core-periphery structure of the system. In order to control for the density of the network, as we increases  $\rho_1$  we have to opportunistically decrease the value of  $\rho_3$ . The average degree  $\langle k \rangle$  can be kept fixed by setting

$$\rho_3 = \frac{2}{(Np) * (Np - 1)} \left( K - K_{cc} - K_{cp} \right). \quad (1)$$

In our case with  $N = 250$  and  $\langle k \rangle = 10$ , we have  $K = 1250$  whereas  $K_{cc}$  and  $K_{cp}$  are set once we fix the core size  $N_c$  and the value of  $\rho_1$ . In Fig. 3 we show the average Jaccard index  $J$  computed for the ground-truth partition and the partition extracted by the algorithm on the stochastic realizations of the network as a function of different values of  $\rho_1$  for different core size.

As shown,  $J$  increases quickly until  $\rho_1 = 0.2$  and only mildly after this point. This indicates that  $\rho_1 = 0.2$ , corresponding to a value of  $\rho_3 = 0.03$ , can be considered as the smallest density of the core-core block at which the core-periphery structure of the network is sufficiently well-defined. For this reason, in the stochastic block model for multiplex networks with different values of core similarity  $S_c$  described in Fig. 1 of the main text, where we have  $N = 250$  and  $N_c = 50$  we set  $\rho_1 = 0.2$ .

Given the set of parameters  $\rho_1$ ,  $\rho_2$  and  $\rho_3$  we can also compute the average degree  $\langle k_c \rangle$  of core nodes

$$\langle k_c \rangle = \rho_1(N_c - 1) + \rho_2(N_p), \quad (2)$$

the average degree  $\langle k_p \rangle$  of the peripheral nodes

$$\langle k_p \rangle = \rho_3(N_p - 1) + \rho_2(N_c). \quad (3)$$

so that we have

$$\langle k \rangle = \frac{N_c \langle k_c \rangle + N_p \langle k_p \rangle}{N}. \quad (4)$$

In Fig. 4 we show the average Jaccard index  $J$  computed for the ground-truth partition and the partition extracted by the algorithm as a function of  $\langle k_c \rangle / \langle k_p \rangle$ . **The Jaccard index  $J$  is defined as**

$$J = \frac{I_c^{[\alpha\beta]}}{N_c^{[\alpha]} + N_c^{[\beta]} - I_c^{[\alpha\beta]}}, \quad (5)$$

where  $N_c^{[\alpha]}$  is the number of core nodes at layer  $\alpha$ ,  $N_c^{[\beta]}$  is the number of core nodes at layer  $\beta$  and  $I_c^{[\alpha\beta]}$  is the number of nodes that are part of the core at both layers  $\alpha$  and  $\beta$ .



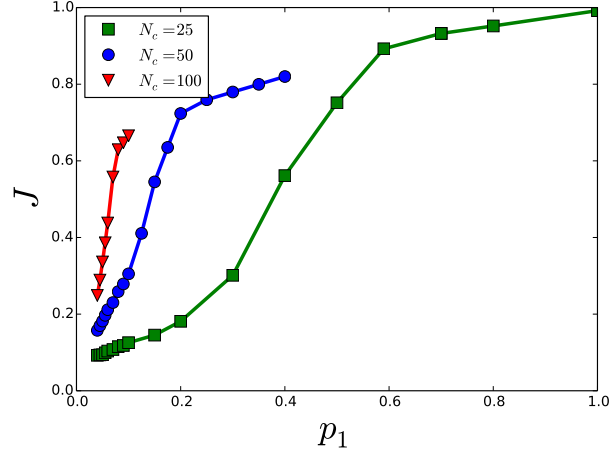


Figure 3: Jaccard index  $J$  for the groundtruth core-periphery partition and the partition obtained by the algorithm on realizations of the stochastic block model as a function of  $\rho_1$  and for different core sizes  $N_c$ .

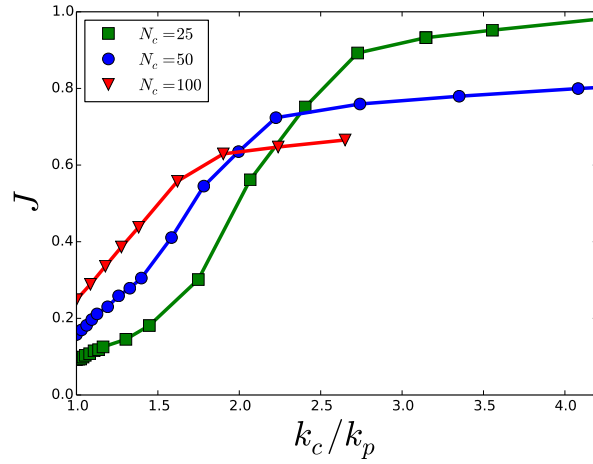


Figure 4: Jaccard index  $J$  for the ground-truth core-periphery partition and the partition obtained by the algorithm on realizations of the stochastic block model as a function of  $\langle k_c \rangle / \langle k_p \rangle$  and for different core sizes  $N_c$ .

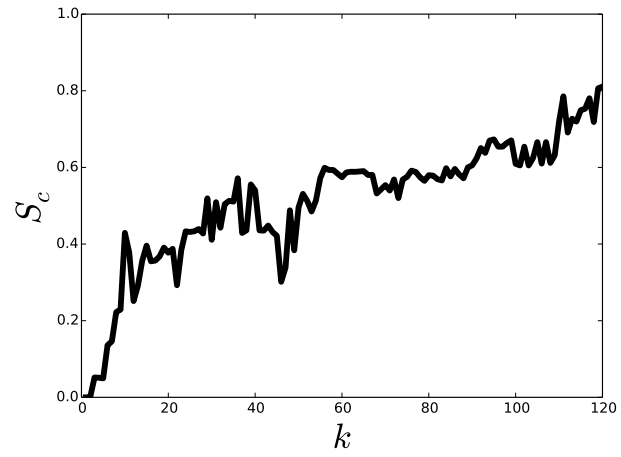
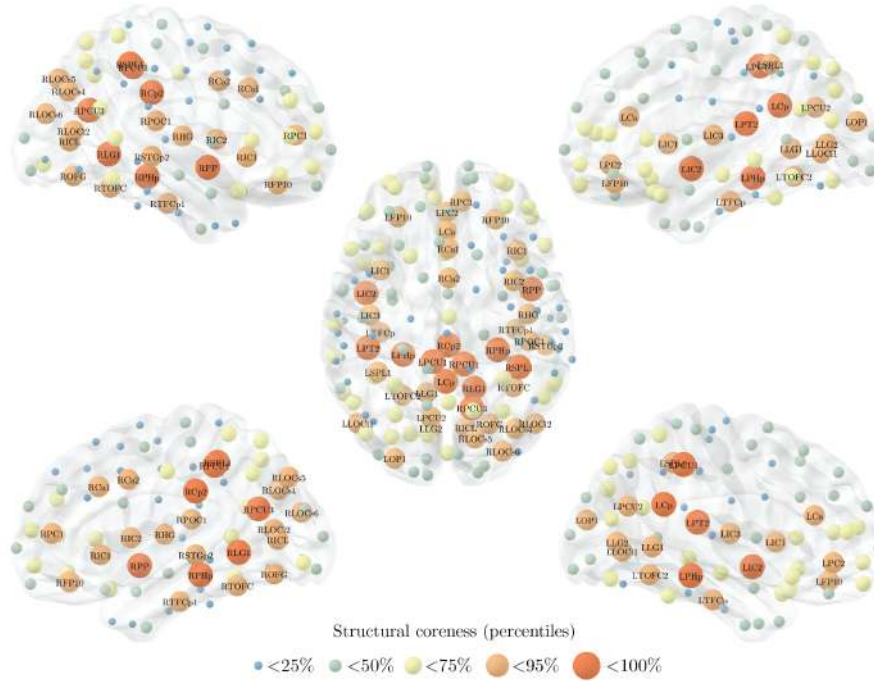


Figure S1: Core similarity  $S_c$  for the structural and functional brain networks thresholded at different values of average degree  $\langle k \rangle$ .

a)



b)

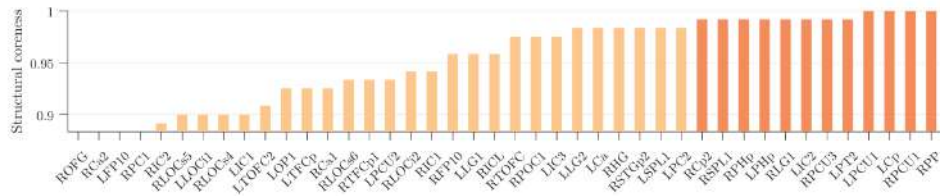


Figure S2: Panel a) shows the node structural coreness from different points of view: external view in the top row, internal view in the bottom row. The color and size of each node code for the percentile to which it belongs as specified in the legend. In panel (b) we report the value of structural coreness of the nodes beyond the 75th percentile with the same color code.

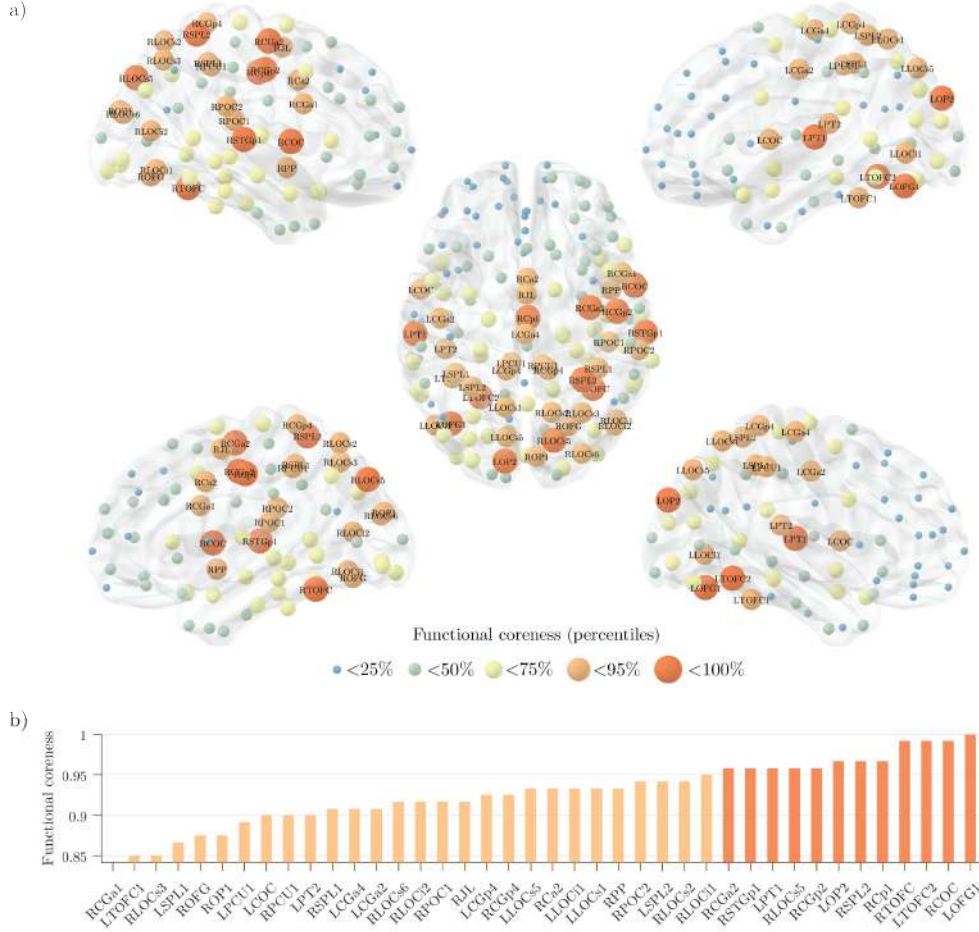


Figure S3: Panel a) shows the functional coreness from different points of view: external view in the top row, internal view in the bottom row. The color and size of each node code for the percentile to which it belongs as specified in the legend. In panel (b) we report the value of functional coreness for the nodes beyond the 75th percentile with the same color code.

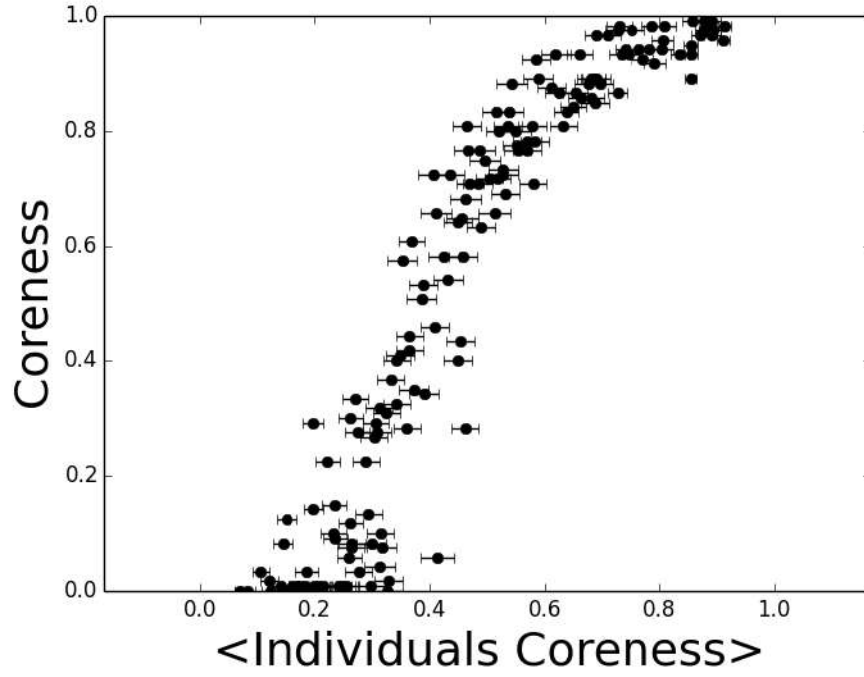


Figure S4: Coreness of the average as a function of average of the coreness. Each point corresponds to a brain region. On the y-axis we show the multiplex coreness of the group-averaged brain networks. On the x-axis we show the average of the multiplex coreness extracted from each individual. Error bars stand for standard error means. The corresponding Spearman's correlation coefficient is  $\rho_s = 0.960$ ,  $p = 2.72 \times 10^{-88}$ .

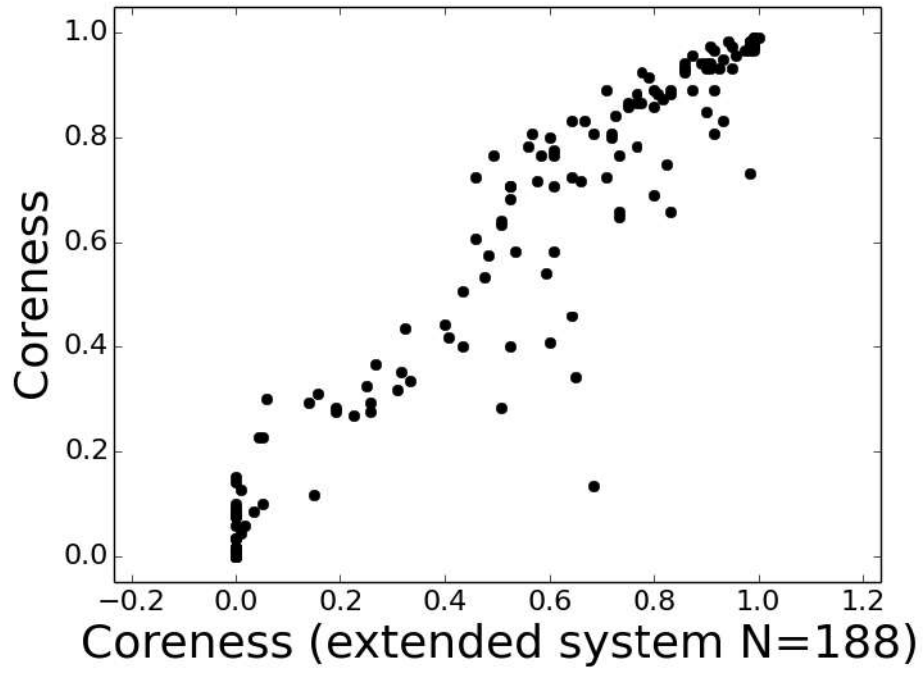


Figure S5: Scatterplot of the multiplex coreness of each ROI computed from brain networks with and without and without subcortical regions. The corresponding Spearman's correlation coefficient is  $\rho_s = 0.95$ , with a  $p = 1.7 \times 10^{-78}$ .





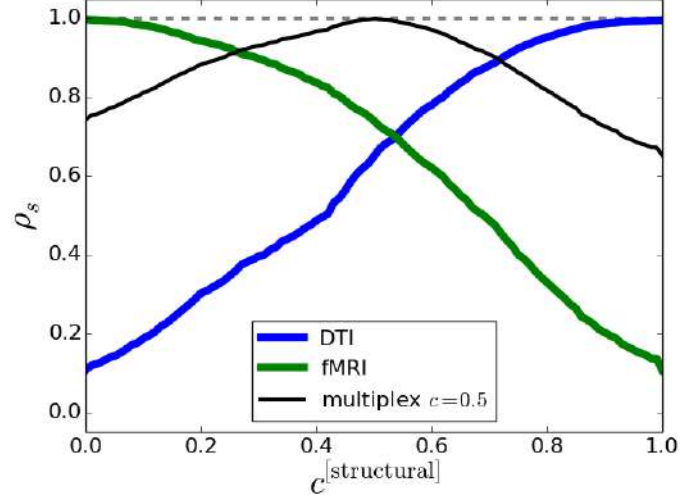


Figure S7: Sensitivity analysis for the multiplex brain coreness. We considered different coefficients  $c^{[\alpha]}$  for the structural and functional layer. Specifically,  $c^{[\text{structural}]} \in [0, 1]$  with  $c^{[\text{functional}]} = 1 - c^{[\text{structural}]}$ . We analyzed the similarity (in terms of Spearman correlation) between the unbiased multiplex coreness and the structural (DTI), functional (fMRI) and multiplex coreness as a function of  $c^{[\text{structural}]}$ . The multiplex coreness is relatively stable across different coefficients around the unbiased case  $c^{[\text{structural}]} = c^{[\text{functional}]} = 0.5$  (black curve); In addition,  $c^{[\text{structural}]} = 0.5$  leads to a multiplex coreness which is slightly more similar to the functional coreness (green curve), highlighting that the multiplex core is more than the sum of the cores at the different layers.

Layer	$N_c$
1	17
2	17
3	12
Multiplex	12

Layer	Layer	$I_C$
1	2	6
1	3	5
2	3	6
Multiplex	1	10
Multiplex	2	8
Multiplex	3	7

Table S1: In the left table we report the size  $N_c$  of the cores of the three layers (mutual trust, common operations, exchanged communications) of the Top Noordin Terrorists network [34] and of the multiplex core shown in Fig. 1 of the main text. In the right table we report the number of common core nodes  $I_c$  belonging to the different pairs of layers. The network is characterized by a core similarity  $S_c = 0.38$  ( $S_c^{[1]} = 0.32$ ,  $S_c^{[2]} = 0.35$ ,  $S_c^{[1]} = 0.46$ . See Eq. 3 in the main text). We also report the number of common core nodes for the multiplex and each layer.

ROI label	Abbrev.	ROI label	Abbrev.
Left Angular	LAG	Right Central Opercular	RCOC
Left Central Opercular	LCOC	Right Cingulate anterior 1	RCa1
Left Cingulate anterior	LCa	Right Cingulate anterior 2	RCa2
Left Cingulate posterior	LCp	Right Cingulate posterior 1	RCp1
Left Frontal Medial	LFMC	Right Cingulate posterior 2	RCp2
Left Frontal Orbital 1	LFOC1	Right Frontal Orbital	RFOC
Left Frontal Orbital 2	LFOC2	Right Frontal Pole 1	RFP1
Left Frontal Pole 1	LFP1	Right Frontal Pole 10	RFP10
Left Frontal Pole 10	LFP10	Right Frontal Pole 2	RFP2
Left Frontal Pole 2	LFP2	Right Frontal Pole 3	RFP3
Left Frontal Pole 3	LFP3	Right Frontal Pole 4	RFP4
Left Frontal Pole 4	LFP4	Right Frontal Pole 5	RFP5
Left Frontal Pole 5	LFP5	Right Frontal Pole 6	RFP6
Left Frontal Pole 6	LFP6	Right Frontal Pole 7	RFP7
Left Frontal Pole 7	LFP7	Right Frontal Pole 8	RFP8
Left Frontal Pole 8	LFP8	Right Frontal Pole 9	RFP9
Left Frontal Pole 9	LFP9	Right Heschls	RHG
Left Inferior Frontal pars triangularis	LIFGpt	Right Inferior Frontal pars triangularis	RIFGpt
Left Inferior Temporal posterior 1	LITGp1	Right Inferior Temporal posterior 1	RITGp1
Left Inferior Temporal posterior 2	LITGp2	Right Inferior Temporal posterior 2	RITGp2
Left Inferior Temporal occipital	LITGt	Right Inferior Temporal occipital	RITGt
Left Insular 1	LIC1	Right Insular 1	RIC1
Left Insular 2	LIC2	Right Insular 2	RIC2
Left Insular 3	LIC3	Right Intracalcarine	RICL
Left Lateral Occipital inferior 1	LLOCi1	Right Juxtapositional Lobule	RJL
Left Lateral Occipital inferior 2	LLOCi2	Right Lateral Occipital inferior 1	RLOCi1
Left Lateral Occipital superior 1	LLOCs1	Right Lateral Occipital inferior 2	RLOCi2
Left Lateral Occipital superior 2	LLOCs2	Right Lateral Occipital inferior 3	RLOCi3
Left Lateral Occipital superior 3	LLOCs3	Right Lateral Occipital superior 1	RLOCs1
Left Lateral Occipital superior 4	LLOCs4	Right Lateral Occipital superior 2	RLOCs2
Left Lateral Occipital superior 5	LLOCs5	Right Lateral Occipital superior 3	RLOCs3
Left Lateral Occipital superior 6	LLOCs6	Right Lateral Occipital superior 4	RLOCs4
Left Lingual 1	LLG1	Right Lateral Occipital superior 5	RLOCs5
Left Lingual 2	LLG2	Right Lateral Occipital superior 6	RLOCs6
Left Middle Frontal 1	LMFG1	Right Lingual 1	RLG1
Left Middle Frontal 2	LMFG2	Right Lingual 2	RLG2
Left Middle Frontal 3	LMFG3	Right Middle Frontal 1	RMFG1
Left Middle Temporal anterior	LMTGa	Right Middle Frontal 2	RMFG2
Left Middle Temporal posterior 1	LMTGp1	Right Middle Frontal 3	RMFG3
Left Middle Temporal posterior 2	LMTGp2	Right Middle Frontal 4	RMFG4
Left Middle Temporal occipital	LMTGt	Right Middle Temporal anterior	RMTGa
Left Occipital Fusiform 1	LOFG1	Right Middle Temporal posterior	RMTGp
Left Occipital Fusiform 2	LOFG2	Right Middle Temporal occipital 1	RMTGt1
Left Occipital Pole 1	LOP1	Right Middle Temporal occipital 2	RMTGt2
Left Occipital Pole 2	LOP2	Right Occipital Fusiform	ROFG
Left Occipital Pole 3	LOP3	Right Occipital Pole 1	ROP1
Left Occipital Pole 4	LOP4	Right Occipital Pole 2	ROP2
Left Paracingulate 1	LPC1	Right Occipital Pole 3	ROP3
Left Paracingulate 2	LPC2	Right Paracingulate 1	RPC1
Left Parahippocampal posterior	LPHp	Right Paracingulate 2	RPC2
Left Parietal Operculum	LPOC	Right Parahippocampal posterior	RPHp
Left Planum Temporale 1	LPT1	Right Parietal Operculum 1	RPOC1
Left Planum Temporale 2	LPT2	Right Parietal Operculum 2	RPOC2
Left Postcentral 1	LCGp1	Right Planum Polare	RPP
Left Postcentral 2	LCGp2	Right Postcentral 1	RCGp1
Left Postcentral 3	LCGp3	Right Postcentral 2	RCGp2
Left Postcentral 4	LCGp4	Right Postcentral 3	RCGp3
Left Precentral 1	LCGa1	Right Postcentral 4	RCGp4
Left Precentral 2	LCGa2	Right Precentral 1	RCGa1
Left Precentral 3	LCGa3	Right Precentral 2	RCGa2
Left Precentral 4	LCGa4	Right Precentral 3	RCGa3
Left Precuneous 1	LPCU1	Right Precuneous 1	RPCU1
Left Precuneous 2	LPCU2	Right Precuneous 2	RPCU2
Left Subcallosal	LSC	Right Precuneous 3	RPCU3
Left Superior Frontal 1	LSFG1	Right Superior Frontal 1	RSFG1
Left Superior Frontal 2	LSFG2	Right Superior Frontal 2	RSFG2
Left Superior Frontal 3	LSFG3	Right Superior Parietal Lobule 1	RSPL1
Left Superior Parietal Lobule 1	LSPL1	Right Superior Parietal Lobule 2	RSPL2
Left Superior Parietal Lobule 2	LSPL2	Right Superior Temporal posterior 1	RSTGp1
Left Supramarginal anterior	LSGa	Right Superior Temporal posterior 2	RSTGp2
Left Supramarginal posterior	LSMp	Right Supramarginal anterior	RSMa
Left Temporal Fusiform anterior	LTFCa	Right Supramarginal posterior	RSGp
Left Temporal Fusiform posterior	LTFCp	Right Temporal Fusiform anterior	RTFCa
Left Temporal Occipital Fusiform 1	LTOFC1	Right Temporal Fusiform posterior 1	RTFCp1
Left Temporal Occipital Fusiform 2	LTOFC2	Right Temporal Fusiform posterior 2	RTFCp2
Left Temporal Pole 1	LTP1	Right Temporal Occipital Fusiform	RTOFC
Left Temporal Pole 2	LTP2	Right Temporal Pole 1	RTP1
Left Temporal Pole 3	LTP3	Right Temporal Pole 2	RTP2
Right Angular	RAG	Right Temporal Pole 3	RTP3

Table S2: Full list of Regions of Interest (ROIs) and abbreviations. Numbers denote the relative position within a macro area, i.e. higher values stand for more posterior ROIs.